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CONDUCTIVE POLYMERIC COMPOSITIONS FOR LITHIUM BATTERIES

This application claims priority rights based on U.S. Provisional Application Serial No. 60/212,231, filed June 16, 2000 and 60/290,864 filed May 14, 2001. The above-identified provisional applications are hereby incorporated by reference.

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INTRODUCTION

1. Technical Field

The present invention relates to novel highly conductive polyanionic

polymers suitable for use in solid polymeric electrolytes in lithium batteries,

especially secondary lithium batteries.

2. Background

Lithium batteries supply energy to a growing number of portable electrochemical devices and are a promising energy source for larger applications

such as electric automobiles. Accordingly, lithium batteries are the subject of intense research and the effort to improve performance continues.

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A major area of interest has been in the field of electrolytes for lithium cells where high conductivity and transport number for lithium ion has been the goal.

Electrolytes are generally prepared by dissolving a highly-conductive salt in a polymer, usually an ether polymer, to make solid polymeric electrolytes (SPE).

Examples of the "salt-in-polymer" approach include the electrolytes disclosed in US Patent No. 5,849,432, US Patent No. 5,824,433 US Patent No. 5,660,947, and US Patent No. 6,235,433.

A "polymer-in-salt" approach has also been attempted. In this approach, chain polymers are added as a dilute component to impart solidity to molten alkali metal salt mixtures of high conductivity (1). Unfortunately, it has been difficult to find simple salts of lithium that are stable and liquid at room temperature. Examples of the polymer-in-salt approach include US Patent No. 5,962,169, US Patent No. 5,855,809, US Patent No. 5,786,110, US Patent No. 5,506,073 and US Patent No. 5,484,670.

Investigations of weakly coordinating anion groups continue to spur the development of new polymeric materials suitable for inclusion into SPE. Fujinami et al. in US Patent No. 6,210,838, disclose a Lewis acid, the weakly coordinating boroxine ring in a polymeric ether chain. Good conductivities are achieved by adding a salt to the polymer. Although the polymer appears to have good mechanical properties, the reported conductivity is too low for commercial applications. Strauss et al. in US Patent No. 6,221,941 disclose weakly coordinating polyfluoroalkoxide anions for applications in electrochemical devices. A highly conductive salt, bis(oxalato)borate, has recently been discovered (German patent No. *DE* 19829030) and its potential as a SPE is being studied. (6).

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The need for conductive polymers continues to spur the development of new materials. Polymeric films which contain weakly coordinating anionic groups are promising candidates as SPE, as they would have good decoupling characteristics and thus high transport number for cations. Batteries and other ionic devices could be made much smaller and lighter by exploiting these films. (2).

Despite continuing discoveries of highly conductive electrolytic salts, and advances in polymerizing these salts, solid polymer electrolytes for lithium batteries are still needed. Especially sought are weakly coordinating anionic materials that can be fabricated into films with high conductivity.

10 3. Relevant literature

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 - S. S. Zhang, Z. Chang, K. Xu and C. A. Angell, "Molecular and Anionic Polymer System with Micro-Decoupled Conductivities", Electrochimica Acta, 45, 12-29 (2000).

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6.. W. Xu, and C. A. Angell, *Electrochem. and Solid State Lett.*, 4, E1 (2001).

5 SUMMARY OF THE INVENTION

It has been discovered that certain anionic groups may be readily bound into chain polymers to make conductive solid polymers. The novel polymers comprise repeat units of weakly coordinating anions in a polyether backbone at separations determined by the number and nature of repeating spacer groups in the polymer chains. The repeating spacer groups also determine the physical characteristics of the polymer including glass transition temperature and mechanical properties such as flexibility, shear strength and solubility.

A method for preparing the subject polymers is provided. In the method the anionic component of certain electrolytic salts comprising a Group III element, preferably orthoborate is modified by chelation with a capping group to make weakly coordinating anionic moieties when incorporated into the polymeric chains. This property makes them suitable for use as solid polymeric electrolytes (SPE) in lithium batteries

In an important aspect of the invention, certain modifications of the

polyanionic polymers, plasticization and cross-linking, e.g., are provided to enhance
the conductivity and optimize certain physical properties of the polymers. The
modified polymers may be formed into films, coatings and extruded into solid forms
for use in electrochemical devices and especially in lithium batteries and rechargeable
lithium batteries.

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BRIEF DESCRIPTION OF THE FIGURES

Figure 1 illustrates temperature dependence of ionic conductivities of PEG-spaced polyMOBs, P(LiOEG_nB), where the length of spacer EG_n is 3 to 23.

Figure 2 illustrates temperature dependence of ionic conductivities of PPG-spaced polyMOBs, P(LiOPG_nB), where the length of spacer PG_n is 7 to 17.

Figure 3 illustrates temperature dependence of ionic conductivities of PEG-spaced polyMMBs, P(LiMEG_nB), where the length of spacer EG_n is 5 and 14.

Figure 4 illustrates temperature dependence of ionic conductivities of LiBH₄

10 crosslinked PEG-spaced polyMOBs, BCLEG_nB, where the length of spacer EG_n is 5

to 14.

Figure 5A illustrates temperature dependence of ionic conductivity of EC-PC (1:1) plasticized P(LiOEG₃B) with different EC-PC content.

Figure 5B illustrates temperature dependence of ionic conductivity of EC-PC (1:1) plasticized P(LiOEG₅B) with different EC-PC content.

Figure 5C illustrates temperature dependence of ionic conductivity of EC-PC (1:1) plasticized P(LiOEG₉B) with different EC-PC content.

Figure 5D illustrates emperature dependence of ionic conductivity of EC-PC (1:1) plasticized P(LiOEG₁₄B) with different EC-PC content.

Figure 6 illustrates temperature dependence of ionic conductivity of plasticized P(LiOEG₅B) by different EC-PC compositions.

Figure 7 illustrates the room temperature conductivity of P(LiOEG₃B) plasticized by different solvents and solvent mixtures with the variation of lithium concentrations. Comparison is also made with the conductivity of simple LiBOB

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solutions in PC. Remembering that our conductivity is entirely due to Li⁺ cations, these results are seen as highly promising.

Figure 8A illustrates temperature dependence of ionic conductivity of EC-PC (1:1 by wt) plasticized LiBH₄-crosslinked P(LiOEG₅B) with different content of plasticizer.

Figure 8B illustrates temperature dependence of ionic conductivity of EC-PC (1:1 by wt) plasticized LiBH₄-crosslinked P(LiOEG₉B) with different content of plasticizer.

Figure 8C illustrates temperature dependence of ionic conductivity of EC-PC (1:1 by wt) plasticized LiBH₄-crosslinked P(LiOEG₁₄B) with different content of plasticizer.

Figure 9 illustrates temperature dependence of ionic conductivities of two gel electrolytes with composition of 20.92 PPMA-8.09 P(LiOEG3B)-35.45 EC-35.44 PC (Example 8), and 21.05 PMMA-7.91 BCLEG5B-35.53 EC-35.52 PC (Example 9), respectively.

Figure 10A illustrates lithium deposition-stripping process and electrochemical oxidation of 80% EC-PC plasticized P(LiOEG₃B) on stainless steel electrode, at a scan rate of 1 mVs⁻¹ at room temperature. SS area = 1.963×10^{-3} cm².

Figure 10B illustrates lithium deposition-stripping process and electrochemical oxidation of 80% EC-PC plasticized P(LiOEG₃B) on nickel electrode, at a scan rate of 1 mVs⁻¹ at room temperature. Ni area = 1.963×10^{-3} cm².

Figure 10C illustrates lithium deposition-stripping process and electrochemical oxidation of 80% EC-PC plasticized P(LiOEG₃B) on aluminum electrode, at a scan rate of 1 mVs⁻¹ at room temperature. Al area = 1.963×10^{-3} cm².

Figure 10D illustrates lithium deposition-stripping process and electrochemical oxidation of 80% EC-PC plasticized P(LiOEG₃B) on copper electrode, at a scan rate of 1 mVs⁻¹ at room temperature. Cu area = 1.963×10^{-3} cm².

5 DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

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It has been discovered that certain anionic groups may be readily bound into chain polymers to make conductive solid polymers. The novel polymers and their formulations are provided wherein the polymeric compositions comprise repeat units of weakly coordinating anions positioned between polymer chains comprising repeating spacer groups.

The weakly coordinating anions will, for the most part, be tetra-coordinated members of the Group III elements, preferably tetra-coordinated boron and most preferably orthoborate, substituted with appropriate chelating groups for providing the desired weakly coordinating characteristics of the resulting anionic moiety. The chelating groups bind two oxygen members of the orthoborate anion, thus leaving two oxygens free for binding into the polymeric chains. Preferred chelating groups are dibasic acid residues, most preferably oxalato, malonato or succinato. Certain other preferred chelating groups 1,2,tetra(trifluoromethyl) ethylenedialato, aryl, phenyl and R-substituted phenyl wherein R is alkyl or halo, SO₂ and silane-co-tetraethylene glycalato[DMSI]. The chelating group may be a bi-dentate group or may be two monodentate groups.

To space the repeating anions in the polymer chain, the repeating spacer groups are chosen to have a length and structure required to achieve the desired separation. Most generally, the spacer groups are polyethers, which may be the same

or different in each occurrence. Certain preferred polyethers are poly(ethylene glycol), (hereinafter termed PEG), or poly(propylene glycol), (hereinafter termed PPG), of different molecular weights. Certain other spacer groups are siloxanes.

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Certain physical characteristics are also determined by choice of repeating spacer groups in the polymer. Glass transition temperature and mechanical properties such as flexibility, shear strength and solubility are affected by the nature of the repeating spacer groups. At room temperature, for example, the PEG-spaced polymers are almost glassy for short spacer units, e.g. when tri(ethylene glycol) (n = 3) is used in the polymerization. They are almost rubbery (very slowly flowing at high temperatures, but rubbery for short time stresses) when PEG200 ($n \approx 5$) is used in the polymerization, and soft or sticky rubbery when PEG400 ($n \approx 9$) and PEG600 ($n \approx 14$) are used in the polymerization. The polymer from PEG1000 ($n \approx 23$) is partly crystallized (the polyether segment) at room temperature. The three PPG-spaced polymers are all highly viscous liquids. The polymers are soluble in various polar solvents, like acetonitrile, acetone and even chloroform. Slow hydrolysis and alcoholysis of the polymers take place on prolonged exposure to water or alcohols. However, the reaction products are benign.

The choice of spacer group also determines the conductivity of the polyanionic polymers. Figure 1 and Figure 2 show the temperature dependence of ionic conductivity of PEG and PPG spaced polyanionic electrolytes measured during steady cooling, before (B) and after (A) chloroform treatment to remove LiBOB. The conductivities of these polymeric forms are strongly dependent on the length of the PEG or PPG spacer between the anionic groups, which can be characterized by the number of ethyleneoxy or propyleneoxy units, n value. The actual separation of

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anions depends on chain conformations. For equal n value, PPG-spaced polymers are distinctly less conducting than PEG-spaced polymers.

A method for preparing the subject polymers is provided. In the method, the anionic component of certain electrolytic salts comprising a Group III element, preferably orthoborate is modified by chelation with a capping group to make weakly coordinating anionic groups. The anionic groups are then reacted with the polymer chains, preferably polyalkylene oxides comprising a terminal reactive group, preferably hydroxyl, under condensations whereby a condensation reaction occurs between the capped anionic group and the reactive group. The weakly coordinating polyanionic polymer and a small molecule result.

In the preferred method, an oxalato-capped orthoboric acid anion, $B(C_2O_4)(OH)_2, \text{ is provided. A poly(aklylene glycol) is also provided. The capped orthoboric acid and the polyalkylene glycol are allowed to react to form a mono-oxalato orthoborate (a polyMOB" having the formula P(LiOEG_nB) or P(LiOPG_nB), where EG represents ethylene glycol, PG represents propylene glycol and$ *n* $represents the number of the spacer repeat units) eliminating water in a condensation polymerization process to provide the polymeric polyanion of whatever cation was used to charge-compensate the anion. Preferably the cation is monovalent and is lithium or sodium. In other instances, a malonato-capped orthoboric acid anion, <math display="block">B(CR_2C_2O_4)(OH)_2, \text{ wherein R is hydrogen or halo, preferably fluoro, is provided. In yet other instances the anion comprises a succinic acid residue of the formula <math display="block">B((CR_2)bC_2O_4)(OH)_2. \text{ In certain preferred instances the anion comprises 1,2-tetra(trifluoromethyl)ethylenedialato(2-)O,O' [OC(CF_3)_2]. This anion is disclosed in Xu, W. and Angell, C.A, Electrochim. And Solid-State Letters, 3 (8) 366-368 (2000)$

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which is hereby incorporated by reference. In certain other instances, the capping group is silane-co-tetraethylene glycalato [DMSI].

The method may be illustrated by the following equations for the preparation of the lithium borate polymer of PEG:.

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$$\bullet$$
 LiOH + HOOCCOOH + B(OH)₃ \longrightarrow LiB(C₂O₄)(OH)₂ + 2H₂O

m LiB(C₂O₄)(OH)₂ + m HO(CH₂CH₂O)_nH
$$\longrightarrow$$

H{O[Li(C₂O₄)]B(OCH₂CH₂)_n}_mOH + 2m H₂O (2)

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Certain by-products of these reactions may be separated from the desired polyanionic polymer by treatment with a suitable solvent such as acetonitrile or chloroform in which the by-products are poorly soluble. The glass transition temperatures, for example, before and after chloroform extraction are given in Table 1.

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Table 1

DTA data for P(LiOEG_nB)s and P(LiOPG_nB)s before and after chloroform treatment to remove dissolved LiBOB

Spacer	n value	Before CHCl ₃ treatment			After CHCl ₃ treatment		
		T_{R} (°C)	T _c (°C)	T ₁ (°C)	T _g (°C)	<i>T</i> _c (°C)	<i>T</i> ₁ (°C)
PEG	3	a	a	a	-0.9		_
	5	-22.9		_	-20.1		
	9	-41.8			-44.3	<u> </u>	
	14	-51.5	-15.4	6.6	-53.3	-15.4	23.1
	23	-54.5	-2.3	20.6	-63.1	-47.2	39.7
PPG	7	-43.3	—	_	-47.0		
	13	-56.9	1-	_	-56.0		T
	17	-63.4			-61.0	1-	_

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a: Not measured.

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It may be seen that most of the polymers exhibit glass transitions in only the studied temperature range between -150 and 100° C. The glass transition temperature ($T_{\rm g}$) decreases with increasing the spacer length for both types of spacers. Although it is not intended that the invention be bound by explanation of this behavior, it is thought that the effect of increasing spacer length is because the shorter spacer polymer has higher lithium ion concentration, raising the cohesive energy via transient crosslinking. Thus at ambient temperature the segmental mobility, and hence the Li⁺ mobility increases with increasing the spacer length.

It may also be seen from Table 1 that the glass transition temperature has decreased after chloroform treatment in nearly every case. Apparently this is because the LiBOB content has been reduced and the number of transient crosslinking sites has decreased. However, the liquidus temperature for $n \approx 14$ and $n \approx 23$ PEG spaced polymers increases relative to that before chloroform treatment. Clearly, therefore, the LiBOB is dissolved preferentially in the polyether chains domains and this has the usual melting point lowering effect. When the salt is removed the melting point goes up again.

The polyanionic polymers of the present invention have one of the formulae:

$$M_b^{+k}[AL]_p^{-q}$$

wherein

AL is a repeat unit in the chain wherein:

A is an anionic group comprising a Group III element.

The anionic groups are preferably orthoborate and are capped with a chemical group that modifies their anionic bonding strength. In certain preferred embodiments wherein the anionic group is an orthoborate, the capping groups bind pairwise to two

oxygens of the orthoborate leaving two oxygens free to bind into the chain polymer units. The capping groups may be a divalent chelate group that binds both oxygens on the Lewis base or may be more than one group, each binding one oxygen.

The capping groups are preferably dibasic acids, most preferably oxalato or malonato groups In certain preferred instances the capping group is 1,2-tetra(trifluoromethyl)ethylenedialato(2-)O,O' [OC(CF₃)₂]. In certain other instances, the capping group is silane-co-tetraethylene glycalato [DMSI].

Certain other preferred capping groups are SO₂, aryl, phenyl and substituted phenyl

10 L is a polymeric chain group chemically linked to A.

and wherein L comprises a determined number

of spacer groups and has the formula:

 $L = (Z)_n$

wherein

Z is a spacer group; and

n is the number of each said spacer groups
and wherein Z is the same or different in each occurrence; and

Z is preferably chosen from the group comprising alkyl, R-

substituted alkyl, alkoxy and R-substituted alkoxy wherein R is selected from the group comprising hydrogen, halo, alkyl, alkoxy, phenyl and substituted phenyl. In other instances, Z is a polysiloxane having the formula Si [(CR₃)₂]-O(CR₂CR₂O)_n wherein n is independently 2 to about 50, preferably 2 to about 20, and R is hydrogen or alkyl.

Z most preferably is a polyether having the formula [(O(CR₂)_aCR₂]_n wherein n is from 2 to about 100, most preferably 2 to about 20, a is zero to about 20 and R is hydrogen, halo, alkyl or R-substituted alkyl wherein R is halo, alkyl or phenyl.

p is a number from about 1 to about 100.

In these preferred embodiments the ether groups may be the same or different in each occurrence.

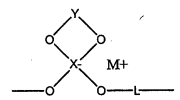
p is the number of repeat units in the polymer.

k is one to 3, most preferably one.

q is one to 3

bq equals bk.

In certain preferred embodiments of the present invention, the repeat group in the polyanionic polymer has one of the formulae:



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wherein X is a Group III element;

O are oxygen;

Y is a capping group.

The counterion M^{+k} is a cation or a cationic group selected from the group comprising hydrogen, Group I metals, Group II metals, NR₄ and PR₄ wherein R is hydrogen, alkyl, or halo, and k is one to three. In

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certain instances wherein the polyanionic polymer is used as an electrolyte in a lithium battery, the counterion is most favorably lithium. In those embodiments wherein the polyanionic polymer is incorporated into an ion exchange system, the cation is preferably a Group I metal or Group II metal, most preferably sodium, potassium, and calcium

It will be appreciated that a molar ratio of a cation to an anion in the polymer of the present invention depends on the valence of the cation and the valence on the anion as well as the number of anionic groups in the polymer. If the total charge on the polymer is k, the ration is reflected in the value of p and k. For example, if both the cation and the anion are monovalent, then k and p are 1, and there will be a 1:1 molar ratio between the cation and the anionic group. Whereas if the cation is divalent and the anion is monovalent, then k is 2 and p is 1, and there will be a 1:2 molar ratio between the cation and the anionic group of the present invention. Preferably, k is an integer from 1 to 3, more preferably 1 to 2, still more preferably k is 1 or 2, and most preferably 1. Preferably p is 1 or 2 and most preferably 1.

The weak charge on the anionic compound acts as a coulombic trap for M^{†p} and as a result, M^{†p} is easily decoupled from the anionic polymer. This decoupling property imparts high cationic conductivity to the polymer and makes the polymers useful as solid polymeric electrolytes in lithium batteries. In certain instances wherein the polyanionic polymer is incorporated into a lithium battery, the counterion is lithium. In those embodiments wherein the polyanionic polymer comprises an ion exchange system, the cation is preferably a Group I metal or Group II metal, most preferably sodium, potassium, calcium.

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Preferred methods for preparing the present polymers comprises providing a precursor anionic group wherein the anionic group comprises a Group III element tetragonally coordinated with oxygen and wherein two of the oxygens are capped with an electron withdrawing group. Also provided is a polymeric group comprising repeating spacer groups and having a reactive group. The anionic group and the polymeric group are combined to form the polyanionic polymer and a small molecule. In preferred embodiments wherein the anionic group comprises a tetra-coordinated oxide of a Group III element wherein two oxygens are capped and two oxygens are coordinated to hydrogen or a cation, the reactive group condenses with the uncapped oxygen and a small molecule such as water or a hydroxide is formed.

The present invention further relates to solid polymeric electrolytes incorporating such polymers and to rechargeable batteries and other electrochemical devices which utilize solid polymeric electrolytes.

Certain modifications can be made to the present conductive polymers to enhance their mechanical properties so they can be more readily formed into films or otherwise fabricated into components suitable for use in secondary lithium batteries. Certain properties of the present polymers indicate their suitability for such purposes. They are soluble in certain solvents and plasticizers, which is a prerequisite for film formation. They may be cross-linked to form polyanionic composites, and these cross-linked composites are likewise soluble or swollen in plasticizers. The polymeric chains in the present polymers may be chosen to be reactive with other polymers so that they may be mixed with, bonded to, or otherwise incorporated into suitable non-ionic chain polymers, ionic chain polymers comprising other ionic groups, polymer networks or block-co-polymers. These modifications have been

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illustrated in the following examples. Certain similar modifications will be apparent to one skilled in the polymer arts.

In an important aspect of the present invention, the polyanionic chain polymers are cross-linked to form a polyanionic polymeric network. Any suitable cross-linking agent may be used, but most preferably the string polymers are chemically crosslinked with lithium boron hydride. Cross-linked polymers exhibit greater mechanical strength than the simple polymer chains.

In yet a further important aspect of the present invention, the polyanionic chain polymers are dissolved in solvents, preferably polar solvents, for example tetrahydrofuran (THF), acetonitrile and acetone. This advantageous property of the polyanionic polymers of the present invention makes them suitable for fabrication into films and coatings.

In a related aspect of the present invention, the polyanionic chain polymers incorporating weakly coordination anionic groups may be affixed to a solid surface and incorporated into an ion-exchange system. The spacer groups may be chosen to provide a tethering group for bonding to a surface such as an ion exchange resin bead or a porous membrane.

In yet another aspect of the present invention, a method is given for increasing the conductivity of the polyanionic polymers wherein certain plasticizers are added to the polymers. Although it is not intended that the present invention be bound by a description of the mechanism of the plasticization effect, it is proposed that the local mobility of the polymeric chain is increased by the plasticizers and as a result the conductivity is increased.

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In an advantageous embodiment of the invention, the anionic chain polymers and the cross-linked network polymers comprise certain plasticizers that enhance the conductivity of the polymer. The plasticized polymers and cross-linked polymers can be formed into conductive films by methods known in the art. Preferred plasticizers are carbonate and non-carbonate plasticizers. Suitable carbonate plasticizers are, for example, ethylene carbonate (EC), propylene carbonate (PC), butylene carbonate (BC), dimethylcarbonate (DMC) and diethyl carbonate (DEC). Suitable non-carbonate plasticizers are 1,2- dimethoxyethane (DME) and 1,2-diethoxyethane (DEE), dimethylsulfoxide (DMSO), dimethylsulfone (DMS), ethylmethylsulfone (EMS), γ-butyrolactone (BL). Preferred plasticizers comprise mixtures of carbonate plasticizer, preferably mixtures of ethylene carbonate and propylene carbonate (EC-PC), ethylene carbonate and dimethyl carbonate (EC-DMC), and propylene carbonate and dimethylxyethane (PC-DME).

The above-mentioned polyanionic polymers and cross-linked polymers and those embodiments wherein the polymers are dissolved in solvents or comprise plasticizers can be employed advantageously as solid polymeric electrolytes in most any type of electrochemical device. Most specifically the polyanionic polymers of the present invention are suitable SPE for electrochemical devices comprising lithium and in particular, lithium rechargeable batteries. The polyanionic polymers can be incorporated in electrochemical cells and lithium batteries, especially rechargeable lithium batteries.

These and other features, aspects and advantages of the present invention will become better understood with reference to the following description, and accompanying drawings.

EXAMPLE 1

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This example illustrates the preparation of a Lewis base-containing polyanionic polymer wherein the Lewis base comprises a tetrakis-coordinated boron structure wherein two ligands are connected with an oxalyl groups and the other two ligands are oxygen linked to the polymeric chain containing repeating spacer units of ethylene glycol. This polymer is generally called polyMOB (wherein "MOB" is for mono-oxalato borate) and has the formula poly[lithium mono-oxalato oligo(ethylene glycol)_n borate] which termed P(LiOEG_nB) wherein n is 3 to 23.

The route to the polyanion polymers is to first form the oxalato-capped orthoboric acid B(C₂O₄)(OH)₂ and then to react this compound with poly(ethylene glycol), PEG, of different molecular weight, eliminating water in a condensation polymerization process to provide the polyanion of whatever cation was used to charge-compensate the anion.

In practice, equal molar quantities of lithium hydroxide monohydrate, oxalic acid dihydrate and boric acid were reacted and all the water was removed by boiling. The solid oxalatoboric acid residue was then refluxed with PEG, chosen from tri(ethylene glycol), PEG200, PEG400, PEG600 and PEG1000, in benzene in a distillation flask equipped with a water separator. The reaction continued until no more water was released. This procedure yielded a rubbery polymer with some fine white particles inside the polymer. The polymer is insoluble in benzene. The supernatant solvent (benzene) was decanted and the residue was washed with fresh benzene once and then evaporated on a rotary evaporator under reduced pressure to obtain a dry gel-like residue. This mass was refluxed with anhydrous acetonitrile to yield a clear solution with white precipitates. The precipitates were filtered off and the

solvent in the filtrate was evaporated thoroughly. Then the residual rubbery solid was dissolved in anhydrous chloroform to yield a clear solution with small amount of a white precipitate. After filtration, the filtrate was evaporated down and the polymer was dried in a vacuum oven at 90°C for 48 hours. The products were rubbery, soft or sticky rubbery, or crystal solid, depending on the PEG used. The ionic conductivities of the PEG-spaced polyMOBs are shown in Figure 1.

Example 2

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This example illustrates the preparation of a Lewis base-containing polyanionic polymer wherein the Lewis base comprises a tetrakis-coordinated boron structure wherein two ligands are connected with an oxalyl groups and the other two ligands are oxygen linked to the polymeric chain containing repeating spacer units of propylene glycol. This polymer is generally called polyMOB (wherein "MOB" is for mono-oxalato borate) and has the formula poly[lithium mono-oxalato oligo(propylene glycol)_n borate] which termed P(LiOPG_nB) wherein n is 7, 13 and 17.

The white product of lithium hydroxide monohydrate, oxalic acid dihydrate and boric acid was refluxed with PPG, chosen from PPG425, PPG725 and PPG1000, in benzene as described in Example 1. The polymers were viscous liquid. The ionic conductivities of the PPG-spaced polyMOBs are shown in Figure 2.

Example 3

This example illustrates the preparation of a Lewis base-containing polyanionic polymer wherein the Lewis base comprises a tetrakis-coordinated boron structure wherein two ligands are connected with an oxalyl groups and the other two ligands are oxygen bound to dimethyl siloxane-co-tetraethylene glycol. This polymer is generally called polyMOB (wherein "MOB" is for mono-oxalato borate) and has

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the formula poly[lithium mono-oxalato oligo(dimethyl siloxane-co-tetraethylene glycolato)_n borate] herein abbreviated as P[LiO(DMSiEG₄)_nB] wherein n is around 12.

To a flame dried 500ml three-neck flask equipped with condenser, thermometer and dropping funnel was added 23.9g (0.123mole) tetraethylene glycol. The flask was heated to 100°C and 18.0g (0.123mole) bis(dimethylamino)dimethyl silane was added dropwise under vigorous stirring. After the addition the reaction was continued at the same temperature while a lot of gas (dimethylamine) was bubbling out of the solution. When the gas evolution nearly ceased (about 2hours), 250ml benzene was added to the reaction flask and followed by adding the product from the reaction of lithium hydroxide monohydrate, oxalic acid dihydrate and boric acid. The azeotropic distillation process was begun and the reaction was treated following the procedures described in Example 1. The polymer was sticky rubbery solid.

This example illustrates the preparation of a Lewis base-containing polyanionic polymer wherein the Lewis base comprises a tetrakis-coordinated boron structure wherein two ligands are connected with an malonyl groups and the other two ligands are oxygen linked to the polymeric chain containing repeating spacer units of ethylene glycol. This polymer is generally called polyMMB (wherein "MMB" is for mono-malonato borate) and has the formula poly[lithium mono-malonato oligo(ethylene glycol)_n borate] which termed P(LiMEG_nB) wherein n is 5 to 23.

The white product from the reaction of lithium hydroxide monohydrate, malonic acid dihydrate and boric acid after evaporating all water was refluxed with PEG, chosen from PEG200, PEG400, PEG600 and PEG1000, in benzene as described

in Example 1. The polymers were rubbery solid. The ionic conductivities of the PEG-spaced polyMMBs are shown in Figure 3.

Example 5

This example illustrates the preparation of a crosslinked polyanionic polymer

from the Lewis base-containing polymers of Example 1. Lithium borohydride

(LiBH₄) was used as the crosslinker.

Polymers from Example 1 (wherein the length of spacer groups was 5 to 14) was dissolved in anhydrous THF and cooled in acetone-dry ice bath. Certain amount of LiBH₄ in THF solution was dropwise added into the above solution with vigorous stirring. After addition, the solution was stirred at room temperature overnight. The solvent was then evaporated at reduced pressure and the residual polymer was dried in a vacuum oven at ca. 70°C for 48 hours. The product was stiff rubber and soft rubber depending on the length of the spacer. The temperature dependence of ionic conductivity of these crosslinked polyanionic electrolytes are shown in Figure 4.

15 Example 6

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This example illustrates the plasticization of a polyanionic polymer prepared in Example 1, in non-aqueous solvents. The non-aqueous solvent is chosen from carbonate, non-carbonate plasticizers or their mixtures. Suitable carbonate plasticizers are, for example, ethylene carbonate (EC), propylene carbonate (PC), butylene carbonate (BC), dimethyl carbonate (DMC) and diethyl carbonate (DEC). Suitable non-carbonate plasticizers are 1,2-dimethoxyethane (DME), 1,2-diethoxyethane (DEE), dimethylsulfoxide (DMSO), dimethyl sulfone (DMS), ethylmethylsulfone (EMS), γ-butyrolactone (BL). Preferred plasticizer mixtures are EC-PC, EC-DMC, EC-DMC-DEC, and PC-DME.

The polyanionic polymer from Example 1 wherein the length of the spacer EG_n was 3 to 14 was mixed well with different amount of plasticizers or plasticizer mixtures. The conductivities of the plasticized electrolytes are given in Figures 5A to 5D, 6 and 7.

5 Example 7

This example illustrates the plasticization of a crosslinked polyanionic polymer prepared in Example 5. The plasticizing effect was measured by using EC-PC(1:1, o/w) as the plasticizer.

The crosslinked polyanionic polymer from Example 5 wherein the length of the spacer EG_n was 5 to 14 was mixed well with different amount of EC-PC (1:1, o/w). The conductivities of the plasticized electrolytes are given in Figures 8A, 8B and 8C.

Example 8

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This example illustrates the preparation of a gel electrolyte containing a Lewis base-containing polyanionic polymer prepared in Example 1 as a polymeric lithium salt.

In a dry glove box, the polyanionic polymer from Example 1 wherein the length of the EG_n spacer was 3 was dissolved in a certain amount of EC-PC (1:1, o/w) mixture in a vial. A quantity poy(methyl methacrylate), PMMA, with high molecular weight of 996,000 was added. The vial was sealed and heated to around 140°C with occasionally shaking till the mixture was well done. The hot viscous mass was pressed in between two stainless steel plates covered with Teflon films. After cooling, the self-standing membrane was pealed off. The conductivity of the gel electrolyte is given in Figure 9.

Example 9

This example illustrates the preparation of a gel electrolyte containing a crosslinked polyanionic polymer prepared in Example 5 as a polymeric lithium salt.

The gel electrolyte was prepared by dissolving PMMA and the crosslinked polyanionic polymer from Example 5 wherein the length of the EG_n spacer was 3 in EC-PC (1:1, o/w) mixture in a vial, as described in Example 8. The conductivity of the gel electrolyte is given in Figure 9.

Example 10

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This example illustrates the electrochemical properties of a plasticized polyanionic polymer prepared in Example 6. The cyclic voltammograms were measured at room temperature on an EG&G potentiostat/galvanostat model 273, with a three-electrode dip-cell with platinum, stainless steel, nickel, aluminum or copper wire as working electrode and lithium metal as counter and reference electrodes. The scan rate was 1 mVs⁻¹. The cyclic voltammetric results are given in Figures 10A to 10

Example 11

D.

This example illustrates the electrochemical properties of a plasticized crosslinked polyanionic polymer prepared in Example 5. The cyclic voltammograms may be measured as described in Example 10.

20 Example 12

This example illustrates the battery performance of an electrolytic solution containing a polyanionic polymer in EC/PC (1:1, o/w) mixture from Example 6, wherein the polyanionic polymer has the spacer length of 3 from Example 1.

Prototype lithium rechargeable batteries were assembled by pressing into appropriate

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cases a sequence of a lithium metal disk anode, a glass fiber film soaked saturatedly with an electrolytic solution of a polyanionic polymer (wherein the length of the polyanionic polymer spacer from Example 1 was 3) in EC/PC (1:1, o/w) mixture from Example 6, and a composite cathode membrane. The latter was a blend of LiCr_{0.015}Mn_{1.985}O₄ as the active intercalation material, carbon black as an electronic conductor and PVdF as a polymer binder, in a weight ratio of 82:10:8. The batteries were assembled in a VAC dry box filled with purified argon. Preliminary investigation into the battery characteristics and performance was performed by examining their galvanostatic charge-discharge cyclic curves.

Those skilled in the art will appreciate that numerous changes and modifications may be made to the preferred embodiments of the invention and that such changes and modifications may be made without departing from the spirit of the invention. It is therefore intended that the appended claims cover all such equivalent variations as fall within the true spirit and scope of the invention.

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We claim:

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2 1. A polyanionic polymer chain having the formula:

- $M_b^{+k}[AL]_p^{-q}$
- 4 wherein
- 5 AL is a repeat unit in the chain wherein:
- A is an anionic group comprising a Group III element;
- 7 L is a polymeric chain group chemically linked to A;
- m is the number of repeat units in the polymer;
- 9 q is the charge on the anion;
- 10 M⁺ is a cation or cationic group;
- P is the number of repeat groups;
- b is the repeat number of cations or cationic groups;
- 13 k is the charge on the cation; and
- bk equals pm.
- 15 L is a chain group comprising a determined number of repeating spacer groups and
- 16 has the formula:
- 17 L = (Z)n
- wherein
- 2 is a spacer group; and
- n is the number of each said spacer groups
- and wherein Z is the same or different in each occurrence.

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- The polyanionic polymer of Claim 1 wherein said anionic group comprises an
 orthoborate group.
- 2 3. A polyanionic polymer chain of Claim 1 wherein AL has the formula:

Q q

4.

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wherein X is a Group III element;

6 O is oxygen;

Y is selected from the group comprising (O)C(CR₂)aCO, wherein a is 0 to 5, SO₂, aryl, phenyl and R-substituted phenyl and R is alkyl or halo 1,3 tetra(trifluromethyl)ethylene dialato, and silane-co-tetraethyleneglycalato; and

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L comprises n spacer groups Z wherein Z is the same or different in each occurrence and is selected from the group comprising alkyl, R-substituted alkyl,, alkoxy and R-substituted alkoxy wherein R is selected from the group comprising hydrogen, halo, alkyl, alkoxy, phenyl and substituted phenyl, and wherein n is 1 to about 30; and p is a number from 1 to 3.

- 2 4. A polyanionic polymer of Claim 1wherein said counterion M^{+k} is a
- 3 cation selected from the group comprising hydrogen, Group I metals, Group II
- 4 metals, NR₄ and PR₄ wherein R is hydrogen, alkyl, or halo; and k is one to three.
- The polyanionic polymer of Claim 1 wherein k is one and M^{+k} is
- 2 lithium, sodium, rubidium or potassium or tetramethylammonium ion.
- 1 6. The polyanionic polymer of f Claim 5 wherein M^{+k} is lithium ion.
- 7. The polyanionic polymer of Claim 3 wherein Y is (O)C(CR₂)aC(O) wherein a
- 2 is 0 to 5 and R is selected from the group comprising hydrogen, alkyl, and halo.
- 1 8. The polyanionic polymer of Claim 3 wherein Y is
- 2 tetra(trifluromethyl)ethylene dialato or silane-co-
- 3 tetraethyleneglycalato.
- 1 9. The polyanionic polymer of Claim 8 wherein Y is oxalato or malonato.
- 1 10. The polyanionic polymer of Claim 3 wherein Z is a polyether having the
- formula [O(CR2)aCR2]n wherein a is zero to about 20, n is from about 2 to 100 and R
- 3 is halo, alkyl or phenyl.
- 1 11. The polyanionic polymer of Claim 3 wherein Z is oligo(ethylene glycol)n or
- 2 oligo(propylene glycol)n wherein n is 2 to about 50.
- 1 12. The polyanionic polymer of Claim 3wherein Z is a polysiloxane having the
- formula Si[(CR3)₂]-O(CR₂CR₂O)_n wherein n is 2 to about 50 and R is hydrogen or
- 3 alkyl.
- 1 13. The polyanionic polymer of Claim 1 wherein said anion A comprises a Group
- 2 III element and oxygen and said polymeric chain group L comprises n spacer groups
- 3 Z selected from the group comprising oligo(ethylene glycol)n or oligo(propylene

4	glycol)n wherein n is 1 to about 30 and wherein Z is the same or different in each				
5	occurrence.				
1	14. The polyanionic polymer of Claim 1 wherein said anion A comprises a borate				
2	anion having two oxygens bound to a dibasic acid residue and two oxygens bound to				
3	polymeric chain groups L, wherein said polymeric chain group L comprises n spacer				
4	groups Z selected from the group comprising [(CR ₂) _a (CR ₂ O)]n, wherein a is zero to				
5	about 50, R is selected from the group comprising hydrogen, halo, hydroxyl, alkyl,				
6	aklenyl, alkoxy, phenyl and substituted phenyl and wherein n is 1 to about 30 and				
7	wherein Z is the same or different in each occurrence.				
1	15. The polyanionic polymer of Claim 1 selected from the group comprising poly				
2	[lithium oxalato oligo(ethylene glycolato)n orthoborate], poly [lithium oxalato				
3	oligo(propylene glycolato)n orthoborate], poly [lithium malonato oligo(ethylene				
4	glycolato)n orthoborate and poly [lithium malonato oligo(propylene glycolato)n				
5	orthoborate wherein n is 3,5,9 or about 14.				
1	16. A method for forming a polyanionic chain polymer of Claim 1 comprising				
2	combining				
3	(a) a weakly coordinating anionic group having the formula:				
4	MA				
5	wherein				
6	A is an anionic group comprising oxygen and a				
7	Group III element and a capping group Y;				
8	M is hydrogen ion, a cation or a cationic group; and				
9	b. a polymeric chain group having the formula:				
10	LB				

wherein B is a reactive group capable of combining with M 11 under conditions whereby L binds to A and the molecule MB is 12 formed. 13 17. The method of Claim 16 wherein 1 A is an orthoborate or an aluminate group 2 Y is bound to two oxygens of said orthoborate or aluminate group and is 3 selected from the group comprising (O)C(CR₂)aCO, wherein a is 0 to 5, SO₂, 4 aryl, phenyl and R-substituted phenyl 1,3 tetra(trifluromethyl)ethylene dialato, 5 and silane-co-tetraethyleneglycalato 6 M is hydrogen or lithium wherein R is alkyl or halo; 7 L comprises (Z)n spacer groups selected from the group comprising (CR₂)_n, 8 and (CR₂O)n wherein n is 1 to about 30, wherein R is hydrogen, alkyl, or halo 9 and wherein Z is the same or different in each occurrence, and 10 B is hydroxyl or halo. 11 The method of Claim 16 wherein said spacer groups are selected from the 1 18. group comprising polyethylene, polypropylene, and isopolypropylene. 2 A polyanionic polymer made by the method of Claim 16, comprising repeat 19. 1 groups having the formula. 2 3

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- 6 X is a Group III element;
- 7 O is oxygen;
- Y is selected from the group comprising comprising (O)C(CR₂)aCO, wherein
- 9 a is 0 to 5, SO₂, aryl, phenyl and R-substituted phenyl and R is alkyl or halo 1,3
- tetra(trifluromethyl)ethylene dialato, and silane-co-tetraethyleneglycalato; and
- 11 L comprises n spacer groups Z wherein Z is the same or different in
- each occurrence and is selected from the group comprising alkyl, R-
- substituted alkyl, alkoxy and R-substituted alkoxy wherein R is
- selected from the group comprising hydrogen, halo, alkyl, alkoxy,
- phenyl and substituted phenyl, and wherein n is 1 to about 30; and
- p is a number from 1 to 3.
- 1 20. The polyanionic polymer of Claim 1 comprising in addition a polymeric
- 2 compound selected from the group comprising a non-ionic chain polymer, a polymer
- 3 comprising cationic or anionic groups, a polymer network or a block co-polymer.
- 1 21. The polyanionic polymer of Claim 1 comprising in addition a salt having high
- 2 conductivity.
- 1 22. The polyanionic polymer of Claim 1 affixed to a solid matrix surface.
- 1 23. The polyanionic polymer of Claim 1 formed into a film.
- 1 24. A cross-linked polyanionic polymer prepared by forming chemical bonds
- 2 between said polymeric chain groups of Claim 1.
- 1 25. A polyanionic polymeric network comprising the cross-linked polymer of 2Claim 24.

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- A polyanionic polymeric network formed by contacting chain polymers of 26. 2
- Claim 1 with boron hydride under conditions whereby cross-linking between said 3
- chains occurs. 4
- The polyanionic polymer of Claim 1 comprising in addition a plasticizer. 27.
- The polyanionic polymer of Claim 27 wherein said plasticizer is a carbonate 28. 1
- plasticizer selected from the group comprising ethylene carbonate, propylene 2
- carbonate, butylene carbonate, dimethyl carbonate, and diethyl carbonate. 3
- The polyanionic polymer of Claim 27 wherein said plasticizer is a non-1 29.
- carbonate plasticizer selected from the group comprising dimethylsulfoxide, dimethyl 2
- sulfone, ethylmethylsulfone, butyrolactone, 1,2-dimethoxyethane, and 1,2-3
- diethoxyethane. 4
- The polyanionic polymer of Claim 27 wherein said plasticizer is a mixture of 30. 1
- carbonate plasticizers selected from the group comprising ethylene carbonate and 2
- propylene carbonate, dimethyl carbonate, and diethyl carbonate. 3
- The polyanionic polymer of Claim 27 wherein said plasticizer is a mixture of 31. 1
- carbonate plasticizers selected from the group comprising ethylene carbonate and 2
- propylene carbonate, dimethyl carbonate and diethyl carbonate. 3
- The polyanionic polymer of Claim 1 in a solvent selected from the group 1 32.
- comprising acetonitrile, acetone, and tetrahydrofuran. 2
- The polyanionic cross-linked chain polymer of Claim 26 comprising in 33. 1
- addition a plasticizer. 2
- A solid polymeric electrolyte comprising a polyanionic polymer of Claim 1. 34. 1
- A solid polymeric electrolyte comprising a polyanionic polymer of Claim 24. 35. 1
- 1 36. A solid polymeric electrolyte comprising a polyanionic polymer of Claim 32.

- 1 37. A solid polymeric electrolyte comprising a polyanionic polymer of Claim 23.
- 1 38. A solid polymeric electrolyte comprising a polyanionic polymer of Claim 20.
- 1 39. The polyanionic polymer of Claim 1 in an electrochemical cell.
- 1 40. The polyanionic polymer of Claim 1 in a lithium battery.

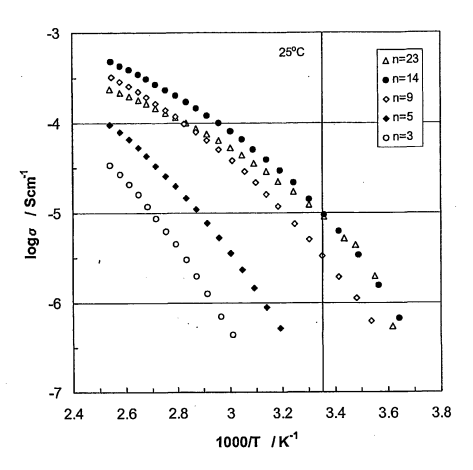


Figure 1. Temperature dependence of ionic conductivities of PEG-spaced polyMOBs, $P(\text{LiOEG}_nB)$, where the length of spacer EG_n is 3 to 23.

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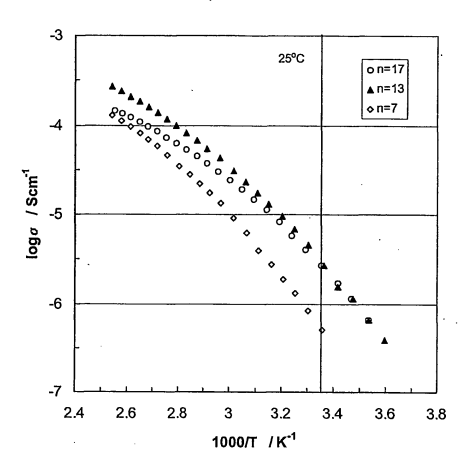


Figure 2. Temperature dependence of ionic conductivities of PPG-spaced polyMOBs, $P(\text{LiOPG}_nB)$, where the length of spacer PG_n is 7 to 17.

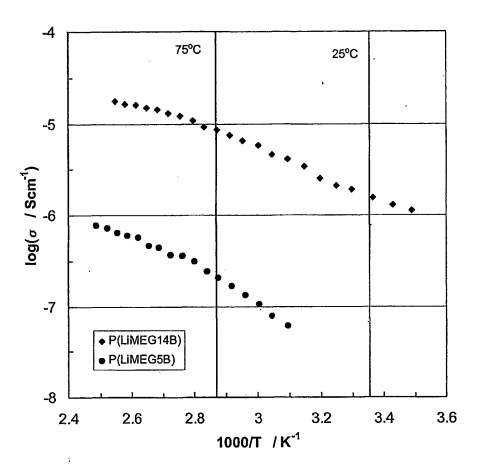


Figure 3. Temperature dependence of ionic conductivities of PEG-spaced polyMMBs, $P(LiMEG_nB)$, where the length of spacer EG_n is 5 and 14.

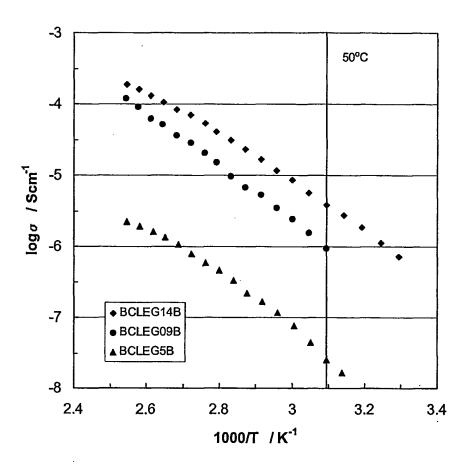


Figure 4. Temperature dependence of ionic conductivities of LiBH₄ crosslinked PEG-spaced polyMOBs, BCLEG_nB, where the length of spacer EG_n is 5 to 14.

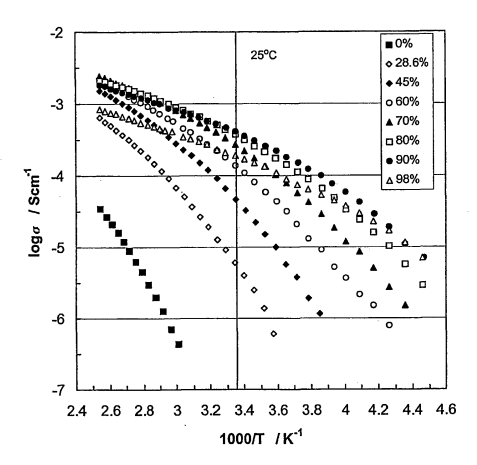


Figure 5A. Temperature dependence of ionic conductivity of EC-PC (1:1) plasticized P(LiOEG₃B) with different EC-PC content.

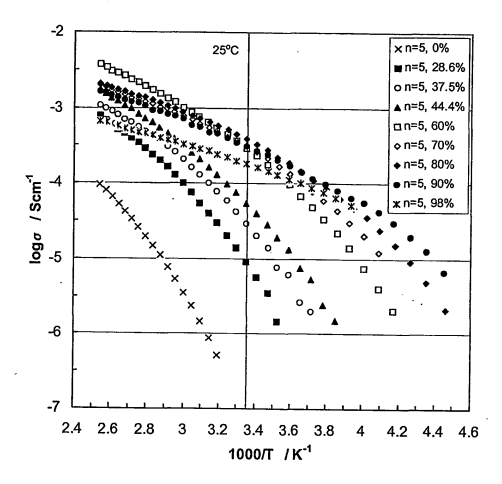


Figure 5B. Temperature dependence of ionic conductivity of EC-PC (1:1) plasticized P(LiOEG₅B) with different EC-PC content.

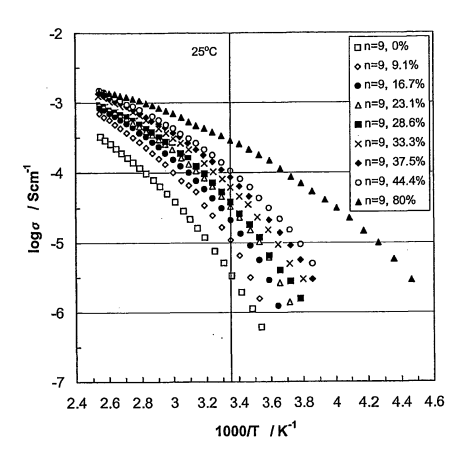


Figure 5C. Temperature dependence of ionic conductivity of EC-PC (1:1) plasticized P(LiOEG₉B) with different EC-PC content.

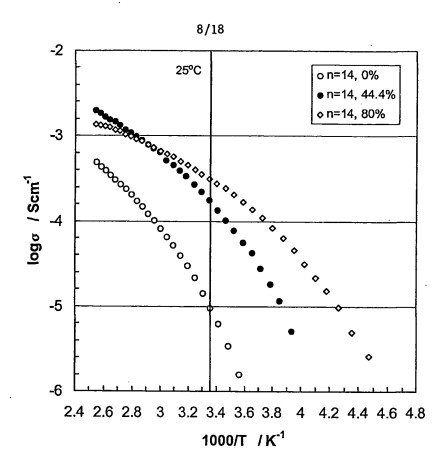


Figure 5D. Temperature dependence of ionic conductivity of EC-PC (1:1) plasticized $P(\text{LiOEG}_{14}B)$ with different EC-PC content.

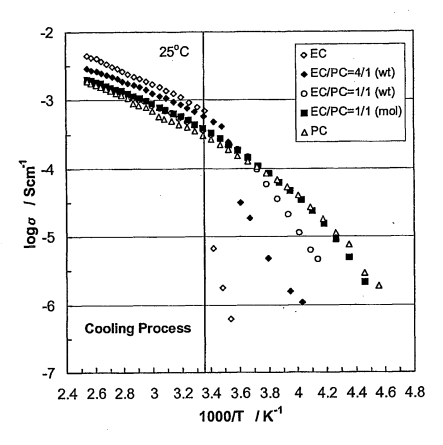


Figure 6. Temperature dependence of ionic conductivity of plasticized $P(LiOEG_5B)$ by different EC-PC compositions.

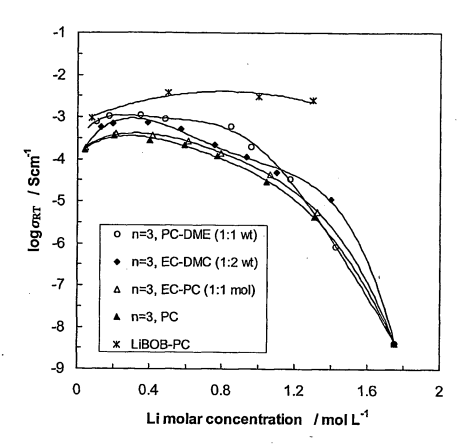


Figure 7. The room temperature conductivity of P(LiOEG₃B) plasticized by different solvents and solvent mixtures with the variation of lithium concentrations. Comparison is also made with the conductivity of simple LiBOB solutions in PC. Remembering that our conductivity is entirely due to Li⁺ cations, these results are seen as highly promising.

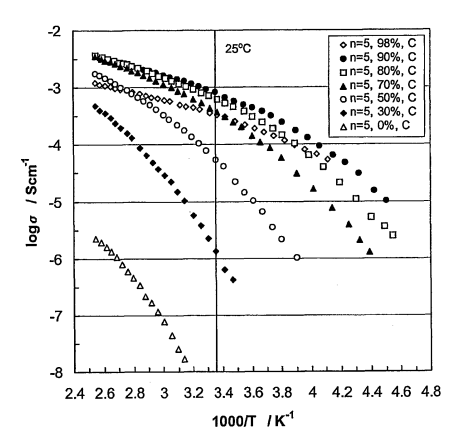


Figure 8A. Temperature dependence of ionic conductivity of EC-PC (1:1 by wt) plasticized LiBH₄-crosslinked P(LiOEG₅B) with different content of plasticizer.

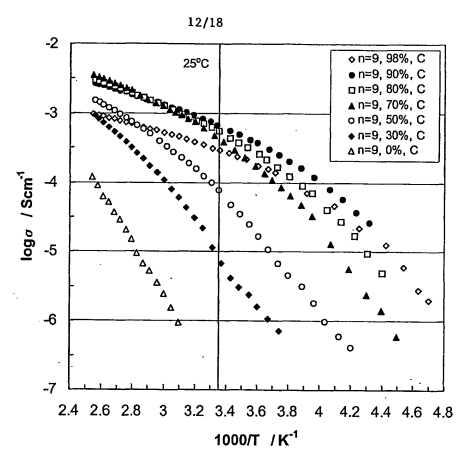


Figure 8B. Temperature dependence of ionic conductivity of EC-PC (1:1 by wt) plasticized LiBH₄-crosslinked P(LiOEG₉B) with different content of plasticizer.

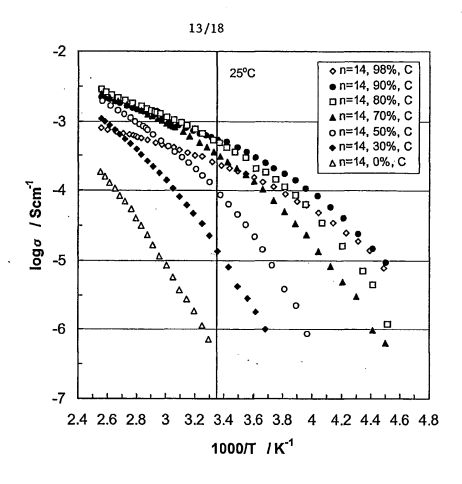


Figure 8C. Temperature dependence of ionic conductivity of EC-PC (1:1 by wt) plasticized LiBH₄-crosslinked P(LiOEG₁₄B) with different content of plasticizer.

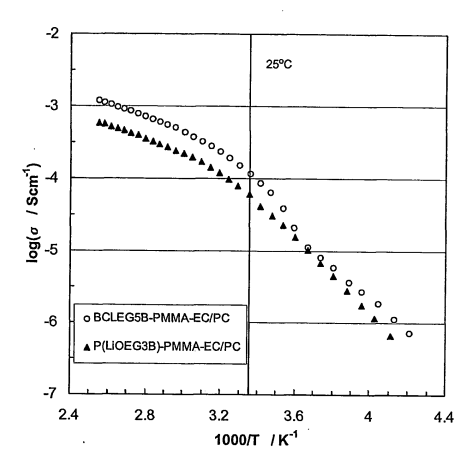
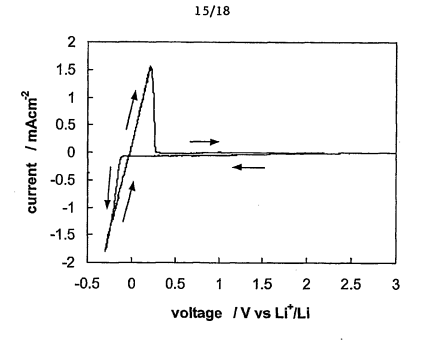


Figure 9. Temperature dependence of ionic conductivities of two gel electrolytes with composition of 20.92 PPMA-8.09 P(LiOEG3B)-35.45 EC-35.44 PC (Example 8), and 21.05 PMMA-7.91 BCLEG5B-35.53 EC-35.52 PC (Example 9), respectively.



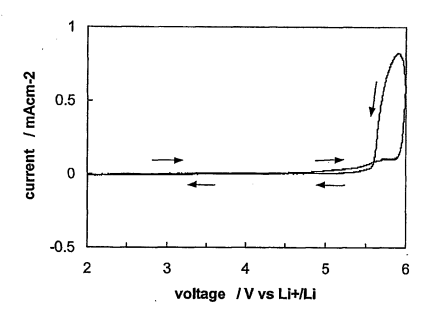
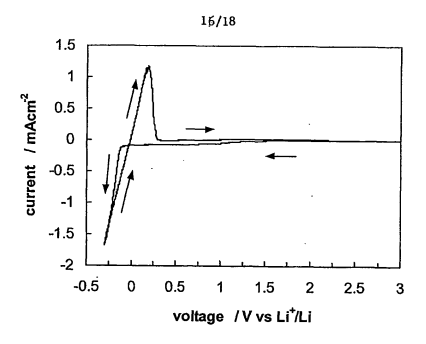


Figure 10A. Lithium deposition-stripping process and electrochemical oxidation of 80% EC-PC plasticized P(LiOEG₃B) on stainless steel electrode, at a scan rate of 1 mVs⁻¹ at room temperature. SS area = 1.963×10^{-3} cm².



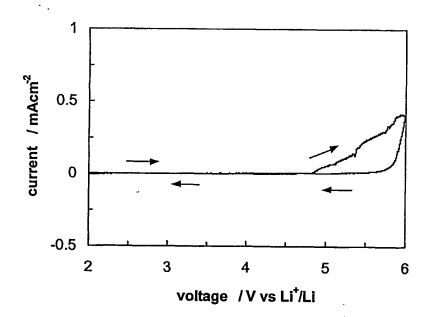
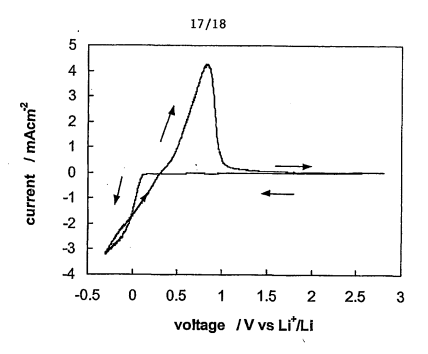


Figure 10B. Lithium deposition-stripping process and electrochemical oxidation of 80% EC-PC plasticized P(LiOEG₃B) on nickel electrode, at a scan rate of 1 mVs⁻¹ at room temperature. Ni area = 1.963×10^{-3} cm².



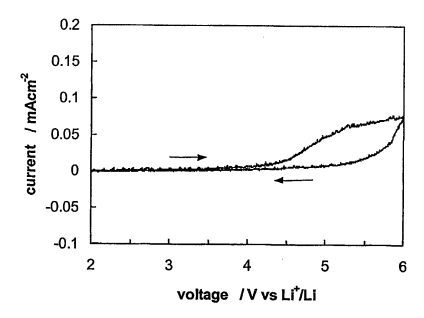
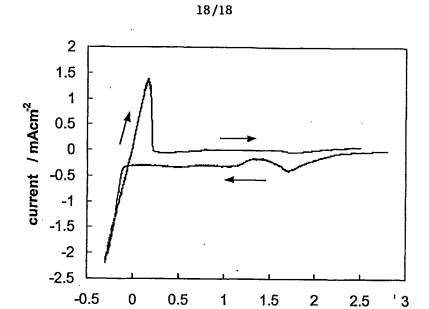


Figure 10C. Lithium deposition-stripping process and electrochemical oxidation of 80% EC-PC plasticized P(LiOEG₃B) on aluminum electrode, at a scan rate of 1 mVs⁻¹ at room temperature. Al area = 1.963×10^{-3} cm².



voltage / V vs Li⁺/Li

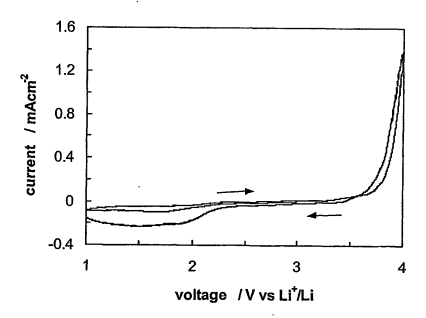


Figure 10D. Lithium deposition-stripping process and electrochemical oxidation of 80% EC-PC plasticized P(LiOEG₃B) on copper electrode, at a scan rate of 1 mVs⁻¹ at room temperature. Cu area = 1.963×10^{-3} cm².

INTERNATIONAL SEARCH REPORT

International application No. PCT/US01/41009

A. CLASSIFICATION OF SUBJECT MATTER						
IPC(7) :CosG 79/08; H01M 10/44, 2/38						
US CL :528/8, 5, 18, 25; 429/50, 51, 192, 209, 212, 245 According to International Patent Classification (IPC) or to both national classification and IPC						
B, FIELDS SEARCHED						
	ocumentation searched (classification system followed	by classification symbols)				
U.S. : 528/8, 5, 13, 25; 429/50, 51, 192, 209, 212, 245						
Documentat	ion searched other than minimum documentation to	the extent that such documents are i	ncluded in the fields			
इन्सिक् रिक्						
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)						
CAS, STN group#	N, search terms: lithium batter?, conductive polymer	P, anionic, (borate or aluminate), weakl	y coordinating anionic			
C. DOCUMENTS CONSIDERED TO BE RELEVANT						
Category*	Citation of document, with indication, where ap	propriate, of the relevant passages	Relevant to claim No.			
Y	ONISHI et al. Thioaluminate polymer complexes as single-ionic solid electrolytes. 1998. Chem Abstract 128: 224296		1-40			
Y	KANBARA et al. Ion-conductive polymand capacitors using it. 04/1996. Chen	1-40				
Y	RAWSKY et al. Aluminosilicate/Poly(ethylene glycol) cvopolymers: A new class of Polyelectrolytes. 1994. Chem Abstract 121: 281368.		1-40			
Y	SAITO et al. Transparent electrically c and laminates thereof, 5/1991. Chem A	1-40				
X Further documents are listed in the continuation of Box C. See patent family annex.						
Special categories of cited documents: "T" later document published after the international filling date or priority						
_	cument defining the general state of the art which is not considered	date and not in conflict with the app the principle or theory underlying th	lication but cited to understand			
to be of particular relevance "X" document of particular relevance; the						
"L" document which may throw doubts on priority claim(s) or which is		considered novel or cannot be conside when the document is taken alone	rea so involve an inventive step			
	ted to establish the publication date of another citation or other ecial reason (as specified)	"Y" document of particular relevance; the considered to involve an inventive step				
	cument referring to an oral disclosure, use, exhibition or other	with one or more other such doom obvious to a person skilled in the art	sents, such combination being			
"P" document published prior to the international filing date but later "&" document member of the same paten than the priority date claimed			family			
Date of the	Date of the actual completion of the international search Date of mailing of the international search report					
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Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT		Authorized officer CEBORAH THOMAS				
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Facsimile N	No. (703) 305-3230	Telephone No. (703) 308-0661				

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US01/41009

C (Continua	tion). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant passages		Relevant to claim No
Y	NOSE et al. Anchoring coatings in electrically conducting plastic films. 3/1991. Chem Abstract 115: 94059		1-40
Y	YAMAGUCHI et al. Copper powder containing electrically conductive coatings. 1/1989. Chem Abstract 112: 8780.		1-40
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